# COMPOSITIONS FOR CLEANING ORGANIC AND PLASMA ETCHED RESIDUES FOR SEMICONDUCTOR DEVICES

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## Abstract of WO 0066697 (A1)

A composition for the cleaning of residues from substrates from about 0.01 percent by weight to about 5 percent by weight of one or more fluoride compounds, from about 20 percent by weight of about 50 percent by weight water, from about 20 percent by weight of about 80 percent by weight of an organic amide solvent and from 0 to about 50 weight percent of an organic suifoxide solvent. The composition has a pH between about 7 and about 10. Additionally, the composition optionally contains corrosion inhibitors, chelating agents, surfactants, acids and bases. In use of the composition, a substrate is contacted with the composition for a time and at a temperature that cermits cleaning of the substrate.

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(54) Title: COMPOSITIONS FOR CLEANING ORGANIC AND PLASMA ETCHED RESIDUES FOR SEMICONDUCTOR DEVICES

#### (57) Abstract

A composition for the cleaning of residues from substrates from about 0.01 persone by weight to about 5 persone by weight to do not one fluoride compounds, from about 20 person by weight to about 50 persone by weight of weight or about 50 persone by weight of about 50 persone by weight of about 50 persone by weight of an organic amide solvent and from 10 to about 50 weight person of on organic sollovide solvent. The composition has p4P between about 7 and about 10. Additionally, the composition optionally contains corrosion inhibitors, chelating agents, surfaceaus, acids and bases. In use of the composition, a substrate is contacted with the composition for a time and or a temperature that permits cleaning of the substrate.

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## COMPOSITIONS FOR CLEANING ORGANIC AND PLASMA ETCHED RESIDUES FOR SEMICONDUCTOR DEVICES

#### 5 Background of the Invention

#### Field of the Invention

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This invention relates to a semiaqueous cleaning composition that is particularly useful for cleaning organic and inorganic compounds or "polymers" (post etch residues) from a semiconductor substrate. As used herein, the term "semiaqueous" refers to a mixture of water and organic solvent. The invention also includes methods of using this composition to clean residues from semiconductor substrates. More particularly, the invention describes a semiaqueous cleaning composition and processes for its use. The solutions are organoammonium compound and amine carboxylate compound free and contain fluoride compounds, water, and solvent and optionally contain corrosion inhibitors, chelating agents, surfactants, acids and bases.

#### Description of Related Art

Fluoride containing chemistries have been used for many years to clean prime silicon wafers (wafers that have not yet undergone ion implantation or device construction) in the semiconductor industry. Normally the fluoride chemistry (usually dilute hydrofluoric acid) is used as the last process step in the sequence called "RCA rinses". The substrate is often contaminated from previous process steps with monolayer amounts of metal, anions and/or organic contaminants or surface residues (particles). These contaminants have been shown to have significant impact on the electrical integrity of simple test device structures and they need to be efficiently cleaned without impairing their integrity. Such cleaning methods could include techniques discussed in the technical literature, for example, Int. Conf. On Solid State Devices and Materials, 1991, pp. 484-486 or Kujime, T. et al., Proc. of the 1996 Semi. Pure Water and Chemicals, pp. 245-256 and Singer, P. Semi. International, p.88, Oct. 1995.

Patents that teach methods for cleaning prime wafers with low pH solutions include US Patent 5,560,857 and 5,645,737: 5,181,985: 5,603,849: 5,705,089.

Using fluoride chemistries (usually HF) as a final RCA cleaning step will cause the silicon wafer surface to be in a hydrophobic state (the surface is covered with Si-H groups) which will repel water. During this cleaning step a certain proportion of the wafer surface is dissolved (removed). Unless the cleaning conditions are carefully monitored (time, temperature, solution composition) the substrates can be damaged, as reported by Rafols, C. et al., J. Electroanalytic Chem. 433, pp. 77-83, 1997. Numerous compositions combine water and organic solvents. The water concentration in these HF solutions is very critical. Silica oxide has an etch rate of 21Å/min (@ 25°C) in HF/water, but in isobutanol the rate was reduced to 2.14Å/min and even lower in acetone (an aprotic solvent) the rate was only 0.12Å/min, as reported at NSF/SRC Eng. Res. Center, Environmentally Benign Semiconductor Manufacturing, Aug. 5-7, 1998, Stanford University.

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After the Front End of Line (FEOL) cleaning process the wafer proceeds to the typical Back End of Line (BEOL) manufacturing process for a semiconductor devices, in which the devices might be dynamic random access memories (DRAMs), static random access memories (SRAMs), logic, electrically programmable read only memories (EPROMs), complementary metal on silicon (CMOS), and the like. Etching fabrication technology using chemical reactions (liquid or plasma) has been used as a method of forming a wiring structure on such semiconductor substrates.

A photoresist film is deposited on the wafer to form a mask, then a substrate design is imaged on the film layer, baked, and the undeveloped image is removed with a photoresist cleaner. The remaining image is then transferred to the underlying material (either a dielectric or metal) with reactive etching gases promoted with plasma energy. The etchant gases selectively attack the unprotected area of the substrate. Liquid etching chemistries, usually containing fluoride chemistries have been used extensively over the years to etch metals (Al) and dielectrics. The fluoride chemistries can be very aggressive and can result in isotropic etching (etching equally in all directions). Isotropic etching effects cannot be tolerated with today's needs for tight critical dimension control, though there have been attempts to control the isotropic etch through statistical process control techniques, as reported by Taylor, D., Solid State Technology, July 1998, p. 119.

The usual plasma etching process involves anisotropic (unidirectional) etching while at the same time the byproducts (composed of photoresist, etching gasses and etched materials) are deposited on the sidewall of etched openings as residues.

A disadvantage of forming this protective sidewall deposit is that it can be very difficult to remove the residue after the etching procedure. If the components in these residues are not removed or neutralized in some manner then the residues will absorb moisture and form acidic species that can corrode the metal structures. The resultant acid corrodes wiring materials to bring about an adverse effect such as an increase in electrical resistance and wire disconnection. Such problems frequently occur, in particular in aluminum and aluminum alloys generally used as wiring material. The wafer substrate in contact with acidic materials, if not controlled, can destroy the metal structures.

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Following completion of the etching operation it is necessary that the resist mask be removed from the protective surface to permit finishing operations. It is desirable to develop an improved cleaning composition to remove the organic polymeric substance from a coated inorganic substrate without corroding, dissolving or dulling the metal circuitry or chemically altering the wafer substrate.

Cleaning compositions used for removing photoresist coatings if not already ashed and other substrates have for the most part been highly flammable, generally hazardous to both humans and the environment, and comprise reactive solvent mixtures exhibiting an undesirable degree of toxicity. Moreover, these cleaning compositions are not only toxic, but their disposal is costly since they might have to be disposed of as a hazardous waste. In addition, these compositions generally have severely limited bath life and, for the most part, are not recyclable or reusable.

Side wall residues have been removed with either acidic organic solvents or alkaline organic solvents. The acidic solvents are generally composed of phenolic compounds or chlorosolvent and/or an aromatic hydrocarbon and/or alkylbenzenesulfonic acids. These formulations generally need to be used at temperatures up to and beyond 100°C. These chemistries normally need to be rinsed with isopropanol.

Dilute hydrofluoric acid solutions can under certain conditions remove the sidewall

30 polymers by aggressively attacking the via sidewall of the dielectric and therefore changing the
dimensions of the device, as taught by Ireland, P., Thin Solid Films, 304, pp. 1-12 (1997), and

possibly the dielectric constant. Previous chemistries that contain HF, nitric acid, water and hydroxylamine are aggressive enough to etch silicon, as taught by U.S. Patent 3,592,773 issued to A. Muller. Recent information also indicates that the dilute HF solutions can be ineffective for cleaning the newer CF<sub>x</sub> etch residues, as taught by K. Ueno et al., "Cleaning of CHF<sub>3</sub> Plasma-Etched SiO<sub>2</sub>/SiN/Cu Via Structures with Dilute Hydrofluoric Acid Solutions," *J. Electrochem. Soc.*, vol. 144, (7) 1997. Contact holes opened on to the TiSi<sub>2</sub> have also been difficult to clean with HF solutions since there appears to be an attack of the underlying TiSi<sub>2</sub> layer. There may also be difficulty with mass transport of the chemicals in the narrow hydrophilic contact holes, as taught by Baklanov, M.R. et al., *Proc. Electrochem. Soc.*, 1998, 97-35, pp. 602-609.

The photoresist around the contact hole of common interlayer dielectrics, TEOS (tetraethylorthosilicate) and boron phosphosilicate glass (BPSG), which are commonly used in ultra large scale integration (ULSI) structures for better conformity of step coverage, is usually removed with HF solutions. It is not uncommon for the HF to also attack the dielectric material. Such attack is not desirable (see Lee, C. and Lee, S, Solid State Electronics, 4, pp. 921-923 (1997)).

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The alkaline organic solvents for post etch residue removal can be composed of amines and/or alkanolamines and/or neutral organic solvents. These formulations generally must be used at temperatures above 100 C. Recently a new class of post etch residue cleaning chemistries has been used to clean these substrates. These chemistries include hydroxylamine, amines, alkanolamines and corrosion inhibitors and generally operate at temperatures 20 to 30 degrees lower.

Recently, fluoride-based chemistries have been used in limited cases to remove post etch residues and to a limited extent, photoresist residues from integrated circuit substrates during BEOL (Back End of Line) processes. Many of the wafer cleaning compositions contain fluoride components, specifically hydrogen fluoride. In addition these compositions might contain strong caustic chemicals (choline-derivatives, tetraalkyl ammonium hydroxide, ammonium hydroxide) such as disclosed in US Patent 5,129,955; US Patent 5,563,119; or US Patent 5,571,447, or might use a two-phase solvent system, which contains one phase with hydrofluoric acid and water while a second phase contains a nonpolar organic solvent (ketones, ethers, alkanes or alkenes) (US Patent 5,603,849). Other formulations include hydroxylamine and ammonium fluoride (US Patent 5,709,756, issued to Ward). Additional examples include quaternary

ammonium salt and fluoride based compositions, as disclosed in published European Application 0662705, and organocarboxylic ammonium salt or amine carboxylate and fluoride based compositions, as disclosed in US Patent 5,630,904.

Some chemistries have also included chelating agents to help remove ionic and anionic contamination from the wafer surface (PCT US98/02794) but chelating agents such as citric acid, gallic acid, and catechol among others, can be aggressive toward the aluminum oxide that covers the Al metal lines. Studies by Ohman and Sjoberg show that the strong complexing ability of citric ions can increase the aluminum oxide solubility and thereby expose the metal to further corrosion, by factors of 166 and 468 at pH 5 and 6 (see Ohman et al., J.Chem. Soc., Dalton Trans. (1983), p. 2513).

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Other methods for cleaning metal and metal oxide residues on wafers include spraying water vapor into the plasma ashing chamber followed by introducing fluorine containing gases (hydrofluoric acid) (US Patent 5,181,985) or a liquid containing hydrofluoric acid, ammonium fluoride and water with a pH between 1.5 to less than 7.

There is also a need to remove particulate residues from the wafer surfaces during the BEOL process. Currently most processes rely on an isopropanol and/or DI water rinse with/or without ultra- or megasonic cleaning. For post chemical mechanical polishing (CMP) cleaning of oxide wafers, normally only DI water is used, but with tungsten wafers a combination of dilute NH<sub>4</sub>OH and dilute HF usually are required.

There are five mechanisms for removing impurities (particles and/or ions) from the wafer surfaces:

- Physical desorption by solvents, which involves replacing a small number of strongly absorbed particles with a large volume of weakly adsorbed solvent (changing the interaction of the surface charges);
- Change the surface charge with either acids or bases, i.e. the Si-OH group can be made positive or protonated with acid or made negative with bases by removing the proton;
- 3. Ion complexion by removing adsorbed metal ions by adding acid (i.e. ion exchange);
- 4. Oxidation or decomposition of impurities, which involves oxidation of metals, organic materials or the surface of slurry particles, will change the chemical forces between the impurities and substrate surface. The chemical reaction can either be through redox chemistry or free radicals;

 Etching the oxide surface, which releases the impurity while dissolving a certain thickness of the substrate surface.

Currently available fluoride-based chemistries can help in items #2 and 5, but the cleaning conditions must be carefully controlled. In many cases, the components of the cleaning compositions are relatively toxic reactive solvent mixtures and thus must be subject to stringent use conditions and require hazardous chemical handling procedures and wearing of safety garments and apparet by users so as to avoid contact with the cleaning compositions. Additionally, because many of the toxic components of such cleaning compositions are highly volatile and subject to high evaporation rates, they require special human and environmental safety precautions to be taken during storage and use of the compositions.

Accordingly, there exists a need to develop improved cleaning compositions to efficiently clean a variety of deposits from a wide variety of substrates. Particularly in the field of integrated circuit fabrication, it should be recognized that the demands for improved cleaning performance with avoidance of attack on the substrates being cleaned are constantly increasing. This means that compositions that were suitable for cleaning less sophisticated integrated circuit substrates may not be able to produce satisfactory results with substrates containing more advanced integrated circuits in the process of fabrication.

These compositions should also be economical, environmental friendly and easy to use.

The present invention teaches such a new and improved cleaning composition and a process for its use. This composition is aqueous, dissolves both organic and inorganic substances, and, when used in the process, is able to clean a variety of substrates. The composition provides a more effective cleaning of the substrates, which means more effective residue removal, which in turn means that higher product yields can be obtained from the substrates being cleaned.

#### SUMMARY OF THE INVENTION

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The novel cleaning compositions of the invention exhibit synergistically enhanced cleaning action and cleaning capabilities at low temperatures that are not possible from the use of the individual components, or the components in combination with other cleaning components, or the components in combination with other cleaning components such as ethoxyethanolamine or alkylamides.

It is a general object of the invention to provide a semiconductor substrate cleaning composition that is effective at low temperatures.

It is a further object of the invention to provide a post etch residue cleaning composition that inhibits redeposition of metal ions.

It is a further object of the invention to form such a cleaning solution that is not flammable.

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It is a further object of the invention to provide such a cleaning solution having low etch rates of silicon oxide.

It is a further object of the invention to provide such a cleaning solution and a process

which removes post etch residues from metal structures.

It is a further object of the invention to provide such a cleaning solution and a process which removes post etch residues from vias.

It is a further object of the invention to provide such a cleaning solution and a process which removes post etch residues from low k dielectrics.

15 These and related objects are attained through the use of the composition and process disclosed herein.

A composition in accordance with this invention is for the cleaning of residues from substrates and is organoammonium and amine carboxylate free. It comprises from about 0.01 percent by weight to about 5 percent by weight of one or more fluoride compounds, from about 20 percent by weight to about 50 percent by weight water, from about 20 percent by weight to about 50 percent by weight water, from about 20 percent by weight to about 80 percent by weight of an organic amide solvent and from 0 to about 50 weight percent of an organic sulfoxide solvent. The composition has a pH between about 7 and about 10. Additionally, the composition optionally contains corrosion inhibitors, chelating agents, surfactants, acids and bases.

A process for cleaning residue from a substrate in accordance with this invention comprises contacting the substrate with an organoammonium and amine carboxylate free composition comprising one or more fluoride compounds, water and an organic amide solvent at a temperature and for a time sufficient to clean the substrate.

#### BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1A is a triangle diagram plotting corrosion results achieved in practice of the invention as a function of composition.

Figures 1B-1C are scanning electron microscope (SEM) photographs of a portion of a semiconductor wafer after treatment with a composition represented in the triangle diagram of Figure 1A.

Figure 2A is a triangle diagram plotting cleaning results achieved in practice of the invention as a function of composition.

Figures 2B-2C are SEM photographs of a portion of a semiconductor wafer after treatment with a composition represented in the triangle diagram of Figure 2A.

Figure 3 is a triangle diagram plotting pH as a function of compositions in accordance with the invention.

Figure 4 is an SEM photograph of a portion of a semiconductor wafer prior to treatment with a composition in accordance with the invention.

Figures 5A-5D are corresponding SEM photographs of portions of a semiconductor substrate after treatment with compositions in accordance with the invention.

#### DETAILED DESCRIPTION OF THE INVENTION

The cleaning composition of this invention is organoammonium and amine carboxylate

free. Several advantages are achievable by being able to eliminate organoammonium and amine
carboxylate compounds from the compositions. Eliminating these compounds reduces cost of
the product. Organoammonium and amine carboxylate compounds are also a class of compound
called phase transfer catalysts. Such compounds under certain solvent conditions can accelerate
undesirable side reactions; i.e. activation of hydroxide or halide reactions (which can corrode
metal surfaces), as reported by "Phase-Transfer Catalysis in Industry," A Practical Guide and
Handbook; Mar. 1991, PTC Interface, Inc. Marietta, GA. The presence of these compounds in
the chemistries can also introduce additional cationic and anionic contamination as raw material
sources.

The cleaning composition contains one or more fluoride compounds. Suitable fluoride

30 compounds are ammonium fluoride, ammonium bifluoride and hydrogen fluoride. The preferred

fluoride compounds are ammonium fluoride and ammonium bifluoride. If hydrogen fluoride is

employed, a buffer is required to bring the pH between about 7 and about 10. The fluoride compounds are desirably present in an amount of from about 0.01 percent by weight to about 5 weight percent, preferably from about 0.05 weight percent to about 5 weight percent. In general, the lower the concentration of the fluoride compound in the composition, the higher the

temperature of use needs to be.

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The cleaning composition contains one or more organic amide solvents. Suitable organic amide solvents are N,N-dimethylacetamide and N,N-dimethylformamide. The preferred organic amide solvent is N,N-dimethylacetamide. The organic amide solvents can be used either singly or as mixtures. The composition optionally contains alkyl sulfoxides such as dimethyl sulfoxide.

The cleaning composition contains water. Typically high-purity deionized water is used.

The composition optionally contains corrosion inhibitors. Suitable corrosion inhibitors include inorganic nitrate salts such as ammonium, potassium, sodium and rubidium nitrate salts, aluminum nitrate and zinc nitrate.

The composition optionally contains chelating agents. Suitable chelating agents are described in commonly assigned U.S. Patent 5,672,577, issued September 30, 1997 to Lee, which is incorporated herein by reference. Preferred chelating agents include catechol, ethylenediaminetetraacetic acid, and citric acid.

The composition optionally contains surfactants. Suitable surfactants include poly(vinyl alcohol), poly(ethyleneimine) and any of the surfactant compositions classified as anionic, cationic, nonionic, amphoteric, and silicone based. Preferred surfactants are poly(vinyl alcohol) and poly(ethyleneimine).

Some combinations of components require the addition of acids and/or bases to adjust the pH to an acceptable value. The acids suitable for use in the present invention are organic or inorganic. The acids can include nitric, sulfuric, phosphoric, hydrochloric acids (though hydrochloric acid can be corrosive to metals) and the organic acids, formic, acetic, propionic, n-butyric, isobutyric, benzoic, ascorbic, gluconic, malic, malonic, oxalic, succinic, tartaric, citric, gallic. The last five organic acids are examples of chelating agents.

General structure for the organic acids:

Concentrations of the acids can vary from about 1 to about 25 wt percent. The important factor is the solubility of the acid and base products with any additional agents in the aqueous solutions.

The caustic components suitable for use to adjust the pH of the cleaning solution can be composed of any common base, i.e. sodium, potassium, magnesium hydroxides, or the like. The major problem is that these bases introduce mobile ions into the final formulation. Mobile ions could destroy computer chips being produced today in the semiconductor industry. Other bases can include choline (a quaternary amine) or ammonium hydroxide.

#### Operation:

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The method of cleaning a substrate using the cleaning compositions of the present invention involves contacting a substrate having residue thereon, particularly organometallic or metal oxide residue, with a cleaning composition of the present invention for a time and at a temperature sufficient to remove the residue. Stirring, agitation, circulation, sonication or other techniques as are known in the art optionally may be used. The substrate is generally immersed in the cleaning composition. The time and temperature are determined based on the particular material being removed from a substrate. Generally, the temperature is in the range of from about ambient or room temperature to 100°C and the contact time is from about 1 to 60 minutes. The preferred temperature and time of contact for this invention is 25 to 45°C from 2 to 60 minutes. Generally the substrate will be rinsed after using the composition. Preferred rinse solutions are isopropagal and DI water.

The compositions of the invention are particularly useful for removing residue from metal and via features.

The compositions of the invention are particularly useful on low-k dielectrics. Low-k dielectrics are known in the art and include fluorinated silicate glass (FSG), hydrido organo siloxane polymer (HOSP), low organic siloxane polymer (LOSP), nanoporous silica (Nanoglass), hydrogen silsesquioxane (HSQ), methyl silsesquioxane (MSQ), divinysiloxane bis(benzocyclobutene) (BCB), silica low-k (SiLK), poly(arylene ether) (PAE, Flare, Parylene), and fluorinated polyimide (FPI).

Examples of cleaning compositions and processes according to the present invention

suitable for removing resist mask or residues from a substrate are set forth in examples below.

Example 1. A group of cleaning chemistries containing the ingredients of ammonium fluoride, water, one or more amides, such as N,N-dimethylacetamide, N,N-dimethylformamide, 1-methyl-2-pyrrolidinone, N,N-dimethylpropionamide, and dimethyl sulfoxide listed in Table 1 were tested with metal wafers which have stacks of TEOS / Ti / TiN /AICu / TiN (from Bottom to Top) etched with Cl<sub>2</sub>/BCl<sub>3</sub> plasma in a commercially available AMT DPS etcher. The resulting metal wafers with residues were cut into small sample pieces, and then the sample pieces were immersed into the chemistry solutions in Table 1 for cleaning for 5 minutes at room temperature. The sample pieces were taken out, rinsed with deionized water and dried with the flow of N<sub>2</sub>. SEM was performed with a Hitachi 4500 FE-SEM for evaluating cleaning and corrosion effects. Residue removal and corrosion effects on metal stack were assessed by visual comparisons and were all ranked on a scale of 1 to 10. The formulations in table 1 are shown in weight percent.

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	Table t					Time	T	-	detail Lin	es	
Formula	NHLF	DI water	DMAC	DMF	DMSO	(Min.)	Clean	Al	Ti	TiN	Oxide
Λ	7	35	64			5	10	10	10	10	10
В		35		64		5	10	10	10	10	10
С	-	35	32	32	T	5	10	10	10	10	10
D	1	35	32		32	5	9	10	10	10	10
Ε	1	35	i	32	32	5	9.5	10	10	10	10

Reaction Temperature: Room Temperature DI water: Deionized water DMAC; N.N-Dimethylacetamide DMSO: Dimethyl sulfoxide DMF: N.N-Dimethylformamide

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Metal Retention for Metal Lines Clean lines: 10 - complete, 1 - not clean at all Al retention: 10 - no change, 1 - Al layer gone Ti retention: 10 - no change, 1 - Ti layer gone TiN retention: 10 - no change, 1 - Til layer gone Oxide retention: 10 - no change, 1 - Oxide layer gone

The results showed that DMF gave the best cleaning performance with no corrosion. However, DMAC and DMSO were chosen for further study based on a consideration of cleaning performance with no corrosion and toxicity considerations.

Example 2. Based on the results in Example 1, N,N-dimethylacetamide and dimethyl sulfoxide were chosen to optimize the cleaning formulations with ammonium fluoride and water. A commercially available LAM TCP9600 etcher with a Cl<sub>2</sub>/BCl<sub>3</sub> plasma was used for etching metal wafers with a stack of Oxide / Ti / TiN /AlCu /TiN (from bottom to top). The resulting metal wafers with residues were cut into small sample pieces, and the sample pieces were immersed into the cleaning solutions in Table 2 for cleaning for 5 minutes at room temperature.

The sample pieces were taken out, rinsed with deionized water and dried with a flow of N<sub>2</sub>. SEM was performed with a Hitachi 4500 FE-SEM for evaluating cleaning and corrosion effects. Residue removal and corrosion effects on the metal stack were assessed by visual comparisons and were all ranked on a scale of 1 to 10. pH values as reported in Figure 3 were measured with a Orion SA520 meter with glass pH electrode. The formulations in table 2 are shown in weight percent.

		Table 2			Time		N	letal Lin	ies	
Formula	NH₄F	DI water	DMAC	DMSO	(Min.)	Clean	Al	Ti	TiN	Oxide
F	i	27	57	15	5	9.5	10	10	10	10
G	1	27	67	5	5	9.5	10	10	10	10
н	1	27	48	24	5	9.5	10	10	10	10
1	1	39	30	30	5	10	10	10	10	10
J	1	79	10	10	5	8	3	10	10	10
K	1	49	40	10	5	9.5	9	10	10	10
L	I	59	20	20	5	6	8	10	10	10
М	ı	49	10	40	5	8	10	10	10	10
N	ı	30	69	0	5	10	10	10	10	10
0	1	29	25	45	5	10	10	10	10	10
P	1	29	35	35	5	10	10	10	10	10
Q	1	29	45	25	5	9	10	10	10	10
R	ı	29	0	70	5	8	10	10	10	10

Reaction Temperature: Room Temperature DI water: Deionized water DMAC: N,N-Dimethylacetamide DMSO: Dimethyl sulfoxide

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Metal Retention for Metal Lines
Clean lines: 10 – complete, 1 – not clean at all
Al retention: 10 – no change, 1 – Al layer gone
Ti retention: 10 – no change, 1 – Ti layer gone
Ti retention: 10 – no change, 1 – TiN layer gone
Oxide retention: 10 – no change, 1 – TiX layer gone
Oxide retention: 10 – no change, 1 – Oxide layer gone

In order to understand the results obtained with the formulations shown in Table 2, the data was plotted in the three triangle diagrams of Figures 1A, 2A and 3, showing respectively, aluminum corrosion as a function of water, DMAC and DMSO concentration in the solutions, residue cleaning performance as a function of water, DMAC and DMSO concentration in the solutions, and pH of the compositions as a function of water, DMAC and DMSO concentration in the solutions. A comparison of Figure 1B shows the substantial corrosion with a composition having a high water content and a low DMAC and DMSO content. Figure IC shows no corrosion with a higher DMAC and DMSO content and lower water content. Figure 2B shows only partial residue cleaning with a composition having a high water content and a low

DMAC and DMSO content. Figure 2C shows complete residue cleaning corrosion with a higher DMAC and DMSO content and lower water content. Figure 3 viewed in combination with Figures 1A and 2A shows the appropriate pH range for avoiding substantial aluminum corrosion and obtaining optimum cleaning results.

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Example 3. One of the compositions (composition N) from Table 2 was chosen to process metal wafers etched with an AMT DPS etcher with Cl<sub>2</sub>/BCl<sub>3</sub> plasma. After the dry etching process, heavy residues were formed on these metal wafers, as shown in the representative sample of Figure 4. The metal wafers with the residues were cut into small sample pieces, and the sample pieces were immersed into the chosen chemistry solution for removing the heavy residues for 3, 5 and 10 minutes at room temperature. Composition N could clean the residues completely at 5 minutes and beyond but could not clean the residues completely at 3 minutes (Figures 5A-5C). The sample pieces were taken out, rinsed with deionized water and dried with a flow of N<sub>2</sub>. SEM was performed with a Hitachi 4500 FE-SEM for evaluating cleaning and corrosion effects.

Photo-removal and residue-removal technology will further evolve to meet the needs of 300-mm wafer processing. The SEZ tool and other new types of equipment are becoming more and more popular for large size wafer processing with the focus shift toward individual wafers rather a 25-wafer lot. The large wafer size increases the importance of having a damage-free resist ash process and residue removal process to prevent the loss of increasingly expensive wafers. Furthermore, fast processing procedure for each single wafer will accumulate to save significant time to overall process. In order to cut the processing time, 0.1 weight percent monoethanolamine (MEA) was added to the chosen formulation. The new formulation with MEA can completely clean the residues at 3 minutes as shown in Figure 5D.

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One skilled in the art will recognize from the foregoing examples that modifications and variations can, and are expected to be made, to the foregoing cleaning solution in accordance with varying conditions inherent in the production process. The embodiments above are given by way of example. For example, the specific discussion of dielectric preservation is indicative of other metal, metal alloy, and polysilicon structures protected by the proposed cleaning

solution and method. The teaching examples do not limit the present invention, which is defined by the following claims.

All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

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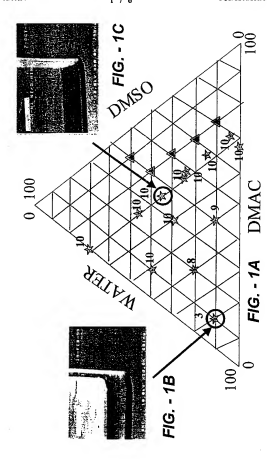
The invention now being fully described, it will be apparent to one of ordinary skill in the art that many changes and modifications can be made thereto without departing from the spirit or scope of the appended claims.

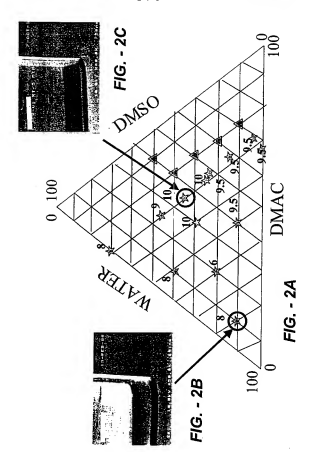
#### WHAT IS CLAIMED IS:

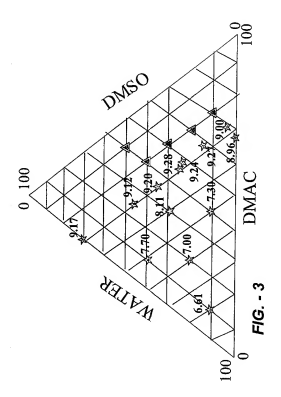
- 1. An organoammonium and amine carboxylate free composition for the cleaning of residues from substrates, comprising from about 0.01 percent by weight to about 5 percent by weight of one or more fluoride compounds, from about 20 percent by weight to about 50 percent by weight water, from about 20 percent by weight to about 80 percent by weight of an organic amide solvent and from 0 to about 50 weight percent of an organic sulfoxide solvent, said composition having a pH between about 7 and about 10.
- 2. The composition of claim 1 further comprising an amine.
- 10 3. The composition of claim 2 in which the amine is an alkanolamine.
  - 4. The composition of claim 3 in which the alkanolamine is monoethanolamine.
  - 5. The composition of Claim 1 further comprising a corrosion inhibitor.
  - 6. The composition of Claim 1 further comprising a chelating agent.
  - 7. The composition of Claim 1 further comprising a surfactant.
- 15 8. The composition of Claim 1 further comprising an acid.
  - 9. The composition of Claim 1 further comprising a base.
  - The composition of Claim 1 comprising ammonium fluoride, water and N,Ndimethylacetamide.
  - 11. The composition of claim 7 further comprising dimethyl sulfoxide.
- 20 12. The composition of Claim 1 comprising ammonium fluoride, water, N,N-dimethylacetamide and an amine.
  - 13. The composition of Claim 1 comprising ammonium fluoride, water, N,N-dimethylacetamide and an alkanolamine.

14. The composition of Claim 1 comprising ammonium fluoride, water, hydroxylamine, alkanolamine, a chelating agent and a corrosion inhibitor.

- 15. A process for cleaning residue from a substrate, which comprises contacting the substrate with an organoammonium and amine carboxylate free composition comprising one or more fluoride compounds, water and an organic amide solvent at a temperature and for a time sufficient to clean the substrate.
- 16. The process of claim 15 in which the temperature is from about 20° to about 100°C.
- 17. The process of Claim 16 where the temperature is about 25° to about 45°C.
- 18. The process of claim 16 in which the time is from about 3 minutes to about 10 minutes.
- 10 19. The process of claim 15 in which the composition comprises from about 0.01 percent by weight to about 5 percent by weight of the one or more fluoride compounds, from about 20 percent by weight to about 50 percent by weight of the water, from about 20 percent by weight to about 80 percent by weight of the organic amide solvent, said composition having a pH between about 7 and about 10.
- 15 20. The process of claim 19 in which the composition additionally comprises up to about 50 weight percent of an organic sulfoxide solvent.
  - 21. The process of claim 20 in which the composition further comprises an amine.
  - 22. The process of claim 21 in which the amine is an alkanolamine.
  - The process of claim 22 in which the alkanolamine is monoethanolamine.







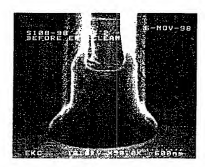


FIG. - 4

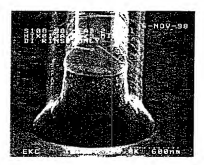


FIG. - 5A

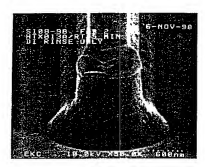


FIG. - 5B

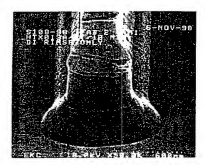


FIG. - 5C

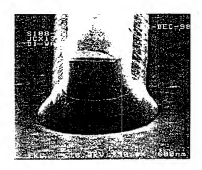


FIG. - 5D

#### INTERNATIONAL SEARCH REPORT

inten nal Application No PCT/US 00/11892

	CI ASSIEL	CATION OF SUBJECT	MATTER		
ĨF	C 7	CATION OF SUBJECT C11D11/00	C1107/50	C11D3/43	G03F7/42

According to International Patent Classification (IPC) or to both national classification and IPC

#### B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 7 C11D G03F

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

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X	EP 0 827 188 A (MITSUBISHI GAS CHEMICAL CO) 4 March 1998 (1998-03-04) example 3; tables 1-1	1,10, 15-19
P,A	WO 99 600° A (ARCH SPECIALTY CHEMICALS INC) 25 N. mber 1999 (1999-11-25) claims 1-25	1-23
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* Special categories of cited documents :	T later document published after the international filing date				
*A* document defining the general state of the art which is not considered to be of particular relevance.	or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention.				
"E" ourlier document but published on or after the international filing date	"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to				
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	imotive an inventive step when the document is taken alons "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the				
"O" document referring to an oral disclosure, use, exhibition or other means	document is combined with one or more other such documents, such combination being obvious to a person skill-				
*P* document published prior to the international fiting date but later than the priority date claimed	in the art. "&" document member of the same patent family				
Date of the actual completion of the international search	Date of mailing of the international search report				
23 August 2000	04/09/2000				
Name and mailing address of the ISA	Ault-orized officer				
European Patent Office. P.B. 5818 Patentiaan 2 NL – 2280 HV Rijswijk					
Tel. (+31-70) 340-2040, Tx, 31 651 epo nl.	Richards, M				

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C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT Category \* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. P,A WO 99 51796 A (OLIN MICROELECTRONIC 1-23 CHEMICALS) 14 October 1999 (1999-10-14) claims 1-18 US 5 709 756 A (PETERS DARRYL W ET AL) Α 1-23 20 January 1998 (1998-01-20) cited in the application claims 1-11 A EP 0 662 705 A (MITSUBISHI GAS CHEMICAL 1-23 CO) 12 July 1995 (1995-07-12) cited in the application example 3 Α EP 0 566 286 A (AMERICAN TELEPHONE & 1-23 TELEGRAPH) 20 October 1993 (1993-10-20) claims 1-6

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